

1 **RESEARH HIGHLIGHT**

2 **The unfolding body plan of primate embryos in culture**

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11 **Improved culture of non-human primate embryos reveals the establishment of the**
12 **crucial framework for subsequent development of bodily tissues and the germline in**
13 **fetuses, bringing us closer to comprehending the elusive development of early human**
14 **embryos.**

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16 Most of what we know about very early mammalian development after blastocyst implants in
17 the womb comes from mouse studies [1]. Little is known about development in primates, and
18 even less so for human development, because of many challenges and constraints on studies
19 on early human embryos. Accordingly, studies on non-human primate embryos might
20 provide information relevant to human development. Two recent studies now report on the
21 development of cynomolgus monkey embryos for 20 days post fertilization, and their
22 progress through gastrulation in culture [2, 3]. They compare their observations on
23 embryonic development in culture by referring to a previous study on the developing monkey
24 embryos in vivo [4].

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26 Shortly after implantation, the inner cell mass (ICM) of humans and non-human primates
27 develop into epithelial cells, followed by the formation of a lumen, which gives rise to the
28 amniotic cavity. The embryonic tissues themselves develop as a bilaminar disc comprising
29 the epiblast on top of the hypoblast; the latter originates from the primitive endoderm. The
30 epiblast and amnion epithelium encapsulate the amniotic cavity while the hypoblast
31 endoderm and extraembryonic mesoderm encapsulate the primary yolk sac. The formation of
32 a primitive streak along the midline of the epiblast disc marks the onset of gastrulation.
33 Gastrulation is a pivotal event during early development when the three primary germ layers,
34 endoderm, ectoderm, and mesoderm, appear for the first time. Primordial germ cells (PGCs),
35 which eventually develop into sperm or eggs, also emerge around this time, either in the
36 amnion according to some studies, epiblast or in both tissues [5-7]. The anterior and posterior
37 ends of the embryos now become distinguishable before the germ layers undergo further
38 differentiation towards organogenesis, in which the anterior cells, e.g., develop into neuronal
39 cells.

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41 In the recent studies on the cynomolgus monkey embryos, Ma et al. [2] and Niu et al. [3]
42 adopted in vitro culture methods previously used to culture mammalian embryos. These
43 studies reveal aspects of non-human primate embryo development after embryo implantation,
44 through gastrulation. These independent studies showed the development of cynomolgus
45 monkey embryos up to 20 days post fertilization (dpf). Niu et al. used the culture system
46 described elsewhere [8]. In contrast, Ma et al. added some novel aspects to their culture
47 conditions containing different serum concentrations, which was first optimized using mouse
48 blastocysts [2].

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50 Broadly speaking, the development of the monkey embryos in both studies recapitulated key
51 developmental events previously described for embryonic development *in vivo* [4, 6]. They
52 observed the formation of the bilaminar disc structure comprising amniotic and yolk sac
53 cavities and the establishment of an anterior-posterior axis. They also saw the emergence of
54 the primitive streak, and the appearance of the PGC-like cells and of the structures
55 resembling neural plate folding [2]. In the future, it will be essential to conduct a rigorous,
56 detailed analysis of development using approaches such as lineage tracing and live imaging
57 to establish the precise origin and destiny of diverse cell types in the embryo, at a time when
58 the embryo undergoes extensive morphogenetic changes.

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60 For now, both studies opted to conduct single-cell transcriptome analysis in the hope to
61 characterize the diverse cell types present in the developing embryos. Ma et al. and Niu et al.
62 compared different stages of embryo development, which they compared with the single-cell
63 transcriptome analysis of cynomolgus monkey embryos *in vivo* described previously [4].
64 Although cells of the majority of embryonic and extraembryonic lineages found *in vivo* were
65 detected in embryos in culture, some disparities in the clustering of cell types were apparently
66 observed. The discrepancies might be due to the appearance of transient and intermediate cell
67 lineages, but the presence of aberrant cells in embryos in culture cannot be excluded.
68 Nonetheless, Ma et al. were able to annotate early (11-14 dpf) and late (16-17 dpf)
69 gastrulating or amniotic cells, where the transcriptional profile of amniotic cells in embryos
70 were not reported before [4]. Niu et al., on the other hand, investigated the chromatin
71 accessibility of the post-implantation embryo at the single-cell level, which, for example, can
72 detect enhancer elements in different cell types during development. Overall, the cell
73 identities from these two studies overlap with those found *in vivo*, suggesting that embryos in
74 culture broadly follow events observed *in vivo*. Note that the culture methods used in these

75 studies can support up to 22% of embryos to the gastrulation stage without fetomaternal
76 interactions. Whether the efficiency of development observed reflects inherent differences in
77 the quality of the embryos or a consequence of culture conditions is unknown. If it is the
78 latter, further optimization of culture conditions, might result in better development at a
79 higher efficiency.

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81 These studies have strengthened our current understanding of the early post-implantation
82 development in primates. They will enhance the technological and conceptual advancement
83 for translating the knowledge to stem cell research and developmental biology. Reliable and
84 efficient culture models could be used to investigate the morphological, molecular, and
85 physiological properties of primate embryos, which could, in principle, be used to study early
86 human peri- to post-implantation development in vitro. There are species-specific differences
87 between early human and non-human primate development, indicating caution when
88 considering extrapolation to human development. The differences include the timing of
89 implantation and amniogenesis between cynomolgous monkey and humans [9], which
90 reflects possible variations due to species evolution [10]. The schedules of gastrulation and
91 establishment of PGCs might also differ, although the regulatory network for PGC
92 specification is apparently broadly conserved in mammals with bilaminar disc embryos [7,
93 10]. Some studies, for example, suggest that PGC-like cells in monkeys can be found in the
94 nascent amnion during the mid-second week (~11 dpf) of development [6]; further work is
95 needed to determine whether amnion and/or epiblast could be the site for the origin of PGCs.
96 Note that PGC specification in the porcine bilaminar disc embryos occurs in the epiblast
97 where the gene regulatory network is more like that in human and not mouse embryos [5].
98 While the developmental events might be broadly similar amongst mammals, there are more
99 significant differences in the development of the extraembryonic tissues. Combined with

possible timing differences amongst mammalian species and their extraembryonic tissues, differences in the molecular mechanism of embryonic development amongst primates cannot be excluded.

The recent studies on gastrulating non-human primate embryos in culture represent a significant advance; much will, however, depend on whether the culture models can be used for more in-depth mechanistic studies. While extrapolation from these studies to early human development is possible, crucial species differences amongst primates cannot be entirely excluded. With legal restrictions on similar studies on human embryos, alternative approaches such as establishing in vitro models for early human development with embryonic stem cells, and generating embryoid-like structures is an option [11]. Increasing improvements and sophistication of the in vitro models might for now address some key questions but cannot substitute entirely for direct studies on very early human development. Such studies, if possible, are of potential value for advances in regenerative medicine and treatment of human diseases.

117 **References**

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Fig. 1 Schematic illustration showing that post-implantation embryo development of primates might be relevant to humans. a Implantation begins with the attachment of blastocyst to the maternal uterine endometrium, when trophoblast cells proliferate into cytotrophoblast and syncytiotrophoblast, which eventually expand and surround the embryo.

b The amniotic cells develop from epiblast cells and form an amniotic cavity, whereas cells from hypoblast form yolk sac endometrium, resulting in a yolk sac cavity. The embryo develops from bilaminar disc epiblast on top of hypoblast cells. Primitive streak appears at the posterior end of embryonic disc, when epiblast starts invading towards hypoblast, thus marking the initiation of gastrulation. Proliferating epiblast cells along the streak migrate to the space between epiblast and hypoblast giving rise to mesoderm cells, thereby converting bilaminar embryonic disc into a trilaminar disc. Primordial germ cells, the precursors of male and female gametes, might originate from the amnion, the epiblast or both tissues during the pre-gastrulation period. They migrate on the wall of the yolk sac close to the allantois, and subsequently along the hindgut before and after colonizing the gonadal ridge. (Figure adapted from Heuser et al., (1941) [12] and Boroviak and Nichols (2017) [13])

